

PATENT COOPERATION RECEAPORTO 14 DEC 2004

PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

REC'D 13 MAY 2004

Applicant's or agent's file reference Case 889/PCT	FOR FURTHER AC	TION See Notifi Prelimina	cation of Fransmittal of International ry Examination Report (Form PCT/IPEA/416)				
International application No. PCT/HU 03/00041	International filing date (d 11.06.2003	lay/month/year)	Priority date (day/month/year) 14.06.2002				
International Patent Classification (IPC) C07D451/04	or both national classification ar	nd IPC					
Applicant SANOFI-SYNTHELABO et al.							
This international preliminary Authority and is transmitted t	examination report has been to the applicant according to A	prepared by this article 36.	International Preliminary Examining				
2. This REPORT consists of a t	2. This REPORT consists of a total of 6 sheets, including this cover sheet.						
heen amended and are	impanied by ANNEXES, i.e. s the basis for this report and ection 607 of the Administration	or sheets contain	cription, claims and/or drawings which have ing rectifications made before this Authority nder the PCT).				
These annexes consist of a t	otal of 4 sheets.						
3. This report contains indication	ns relating to the following ite	ems:					
! ⊠ Basis of the opini	on						
II 🔲 Priority		1					
III 🖾 Non-establishme	nt of opinion with regard to no	ovelty, inventive s	tep and industrial applicability				
IV 🔲 Lack of unity of ir							
V 🖾 Reasoned staten	nent under Rule 66.2(a)(ii) wit lanations supporting such sta	th regard to nove tement	Ity, inventive step or industrial applicability;				
- VI Certain documer		. Andrew Carlot Andrew Carlot Car					
	the international application						
1	ions on the international appli						
Date of submission of the demand		Date of completion	n of this report				
11.12.2003		12.05.2004					
Name and mailing address of the Interpreliminary examining authority:		Authorized Office	gertitude Petanian,				
European Patent Office D-80298 Munich		Steendijk, M					
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 Basis of the r 	eport
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1. With regard to the **elements** of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)):

	De	scription, Pages					
	1-1	3, 15, 16, 18, 20, 22-	as originally filed				
	14,	17, 19, 21	received on 29.04.2004 with letter of 26.04.2004				
	Cla	ims, Numbers					
	1-2	0	as originally filed				
	Dra	wings, Sheets					
	1/4		as originally filed				
With regard to the language, all the elements marked above were available or furnished to this Autholanguage in which the international application was filed, unless otherwise indicated under this item.							
	The	ese elements were av	ailable or furnished to this Authority in the following language: , which is:				
		the language of a tra	anslation furnished for the purposes of the international search (under Rule 23.1(b)).				
		the language of a translation furnished for the purposes of international preliminary examination (under Rule 55.2 and/or 55.3).					
3.	Witl inte	n regard to any nucle mational preliminary (otide and/or amino acid sequence disclosed in the international application, the examination was carried out on the basis of the sequence listing:				
		contained in the international application in written form.					
		filed together with the	e international application in computer readable form.				
		furnished subsequer	ntly to this Authority in written form.				
	☐ furnished subsequently to this Authority in computer readable form.						
		The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.					
		The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.					
4.	The	esulted in the cancellation of:					
		the description,	pages:				
		the claims,	Nos.:				
		the drawings,	sheets:				

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5.		This report has been establis been considered to go beyon	hed as d the d	s if (some of) disclosure as	the amendments had not been made, since they have filed (Rule 70.2(c)).		
		(Any replacement sheet contreport.)	aining	such amend	ments must be referred to under item 1 and annexed to the		
6.	Add	litional observations, if necess	ary:				
111.	. Nor	n-establishment of opinion w	ith re	gard to nove	elty, inventive step and industrial applicability		
1.	The obv	questions whether the claime ious), or to be industrially appl	d inve	ntion appear have not bee	s to be novel, to involve an inventive step (to be non- en examined in respect of:		
		the entire international application,					
	\boxtimes	claims Nos. 14-20					
		because:					
	⊠	the said international application, or the said claims Nos. 14 relate to the following subject matter which does not require an international preliminary examination (specify):					
		see separate sheet					
		the description, claims or drawings (indicate particular elements below) or said claims Nos. are so unclear that no meaningful opinion could be formed (specify):					
		the claims, or said claims Noscould be formed.	. are s	o inadequate	ely supported by the description that no meaningful opinion		
		no international search report	has b	een establish	ned for the said claims Nos. 15-20		
2.	or a	meaningful international preliminary examination cannot be carried out due to the failure of the nucleotide and amino acid sequence listing to comply with the standard provided for in Annex C of the Administrative structions:					
		the written form has not been	furnisl	ned or does i	not comply with the Standard.		
		the computer readable form h	as not	been furnish	ned or does not comply with the Standard.		
 V.	Rea	soned statement under Artic	le 35(2) with rega	rd to novelty, inventive step or industrial applicability;		
	cita	tions and explanations supp	orting	such state	ment		
1.	Stat	ement					
	Nov	eity (N)	Yes: No:	Claims Claims	1-14		
	Inve	ntive step (IS)	Yes: No:	Claims Claims	1-14		
	Indu	strial applicability (IA)	Yes: No:	Claims Claims	1-13		

2. Citations and explanations

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see separate sheet

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INTERNATIONAL PRELIMINARY . International application No. PCT/HU03/00041 EXAMINATION REPORT - SEPARATE SHEET

1) The present application relates to DPP-IV inhibitors of formula I (claims 1-14) as well as intermediates comprising either the azabicyclic ring B or the proline analogue Z (claims 15-20).

No additional search fees were paid for claims 15-20 following an invitation in accordance with R 40 PCT. No preliminary examination can be carried out for the matter not covered by the search report.

2) The following documents are cited herein:

D1: WO-A-98 19998

D2: WO-A-01 34594

D3: EP-A-1 323 710

D4: WO-A-01 96295

D5: US-A-4 273 778

D6: WO-A-03 02553

Document D6 was published after the claimed priority date; on the presumption that the priority is valid, this document is not regarded as prior art.

3) Novelty (claims 1-14)

Documents D1-D4 describe DDP-IV inhibitors derivatives which differ from the presently claimed compounds of formula I in the presence of a different group at the position of the azabicyclic ring B.

Document D5 describes intermediates comprising an azabicyclic-amine, which lack the specific proline-like group Z.

It is noted that document D6 describes DDP-IV inhibitors derivatives comprising an azabicyclic ring; the compounds of D6 differ, however, in the further subsitutions.

4) Inventive step (claims 1-14)

Documents D1-D4 may be considered to represent the closest prior art; the structurally nearest compounds are those carrying a piperidine in stead of an azabicyclic ring for B.

The applicant has provided comparative date with respect to compound of the closest prior art indicating that the bicyclic analogues presently claimed provide for

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INTERNATIONAL PRELIMINARY International application No. PCT/HU03/00041 EXAMINATION REPORT - SEPARATE SHEET

a particular high level of activity (IC50 below 20 nM in Caco-2 test), which would seem the more surprising as the tested pyrolidine/piperidine analogs tend to lower activities than the open chain derivatives.

Accordingly, as solution to the problem of providing further and improved DDP-IV inhibitors, the claimed subject-matter may not be considered obvious.

5) Further observations

Claim 14 relates to subject-matter considered by this Authority to be covered by the provisions of Rule 67.1(iv) PCT. Consequently, no opinion will be formulated with respect to the industrial applicability of the subject-matter of this claim (Article 34(4)(a)(i) PCT).

The claims rely on the drawings for the definition of the subject-matter; the claims should, however, be clear per se.

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Example 1.

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(4R)-3-(2-{[8-(2-Pyrimidinyl)-8-azabicyclo[3.2.1]oct-3-yl]exo-amino}acetyl) thiazolidine-4-carbonitrile

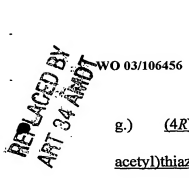
The meaning of R is 2-pyrimidinyl group, B means a group of formula (1), Z means a group of formula (A) in general formula (I).

a.) <u>tert-Butyl 8-(2-pyrimidinyl)-8-azabicyclo[3.2.1]oct-3-yl-exo-carbamate</u> with the general formula (V) - where R and B are given above, Y is tert-butoxycarbonyl group

14,7 g (65 mmol) of *tert*-butyl 8-benzyl-8-azabicyclo[3.2.1]oct-3-yl-*exo*-carbamate (*J. Med. Chem.* **1991,** 34, 656) and 8,93 g (78 mmol) of 2-chloropyrimidine and 12,7 ml (85 mmol) of 1,8-diazabicyclo[5.4.0]undec-7-ene were dissolved in 230 ml of *n*-pentanol and heated under reflux for 4 hours. The solvents were evaporated and the residue was dissolved in 250 ml of chloroform and washed with 2x300 ml of water, dried over sodium sulfate, and purified by column chromatography using *n*-hexane - ethyl acetate- chloroform (1:1:1) as eluent to result in white crystals which were triturated with *n*-hexane. Yield: 13,25 g (67%). M.p.: 113-115°C. ¹H-NMR (CDCl₃): δ 1.34 (s, 9H), 1.49 (t, 2H), 1.66-1.97 (m, 6H), 3.89 (br, 1H), 4.61 (d, 2H), 6.60 (t+br, 1+1H), 8.34 (d, 2H).

b.) 8-(2-Pyrimidinyl)-8-azabicyclo[3.2.1]oct-3-yl-exo-amine with the general formula (II), where R and B are given in step 1a.)

13 g (43 mmol) of *tert*-butyl 8-(2-pyrimidinyl)-8-azabicyclo[3.2.1]oct-3-yl-*exo*-carbamate was dissolved in a mixture of 120 ml of trifluoroacetic acid and 120 ml of dichloromethane. The solution was stirred for 30 minutes and evaporated. The residue was dissolved in 50 ml of dichloromethane and evaporated. This method was repetead three



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g.) (4R)-3-(2-{[8-(2-Pyrimidinyl)-8-azabicyclo[3.2.1]oct-3-yl]exo-amino} acetyl)thiazolidine-4-carbonitrile

245 mg (1,2 mmol) of 8-(2-pyrimidinyl)-8-azabicyclo[3.2.1]oct-3-yl-exo-amine and 191 mg (1 mmol) of (4R)-3-(2-chloroacetyl)thiazolidine-4-carbonitrile and 0,42 ml (3 mmol) of triethylamine were dissolved in 20 ml of dry acetonitrile and stirred at 70°C for 4 hours and then at room temperature overnight. Then the mixture was evaporated to give a yellow thick oil which was purified by column chromatography using chloroform-methanol (9:1) as the eluent to result in a solid white product which was crystallized from diethyl ether. Yield: 191 mg (53%). M.p.: 135-136°C. ¹H-NMR (400 MHz, DMSO-d₆): δ 1.33 (td, 2H), 1.6-2.0 (m, 5H), 3.05 (tt, 1H), 3.32 (m, 2H), 3.44 (ddd, 2H), 4.63 (s, 2H), 4.56 (d, 1H), 4.61 (m,2H), 4.70 (m,1H), 5.23 (dd, 1H), 6.60 (t, 1H), 8.33 (m, 2H).

Example 2.

(4R)-3-(2-{[8-(5-Cyanopyridin-2-yl)-8-azabicyclo[3.2.1]oct-3-yl]-exo-amino} acetyl)thiazolidine-4-carbonitrile dihydrochloride

In the general formula (I) R stands for 5-cyanopyridin-2-yl group, B means for the group of formula (1), Z stands for the group of formula (A).

a.) <u>tert-Butyl 8-(5-cyanopyridin-2-yl)-8-azabicyclo[3.2.1]oct-3-yl-exo-carbamate</u> with general formula (V), where R and B are given above, Y is tert-butoxycarbonyl group

The solution of 415 mg (3 mmol) of 2-chloro-5-cyanopyridine, 679 mg (3 mmol) of tert-butyl 8-benzyl-8-azabicyclo[3.2.1]oct-3-yl-exo-carbamate and 0,46 ml (3,1 mmol) of diazabicyclo[5.4.0]undecene in 25 ml of n-pentanol was refluxed for 8 hours. The resulting solution was evaporated in vacuum, the residue was dissolved in dichloromethane, washed with water and dried over sodium sulfate. After purification by chromatography using n-





precipitation with diethyl ether the title compound was obtained in the form of white crystals: 75 mg (32 %), mp: 204-206°C. ¹H-NMR (DMSO-d₆): δ 1.70-1.78 (m, 4H), 2.01 (m, 4H), 3.37 (m, 2H), 3.67 (m, 1H), 4.07 (m, 1H), 4.21 (m, 1H), 4.56 (d, 1H), 4.76-4.79 (m, 3H), 5.33 (m, 1H), 6.89 (d, 1H), 7.91 (dd, 1H), 8.53 (d, 1H), 9.01 (bs, 2H).

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Example 3.

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(4R)-3-(2-{[8-(2-Pyrazinyl)-8-azabicyclo[3.2.1]oct-3-yl]exo-amino}acetyl) thiazolidine-4-carbonitrile dihydrochloride

The meaning of R is 2-pyrazinyl group, B means a group of formula (1), Z means a group of formula (A) in general formula (I).

a.) <u>tert-Butyl 8-(2-pyrazinyl)-8-azabicyclo[3.2.1]oct-3-yl-exo-carbamate</u>
with the general formula (V) - where R and B are given above, Y is tert-butoxycarbonyl
group

0,54 ml (6 mmol) of chloropyrazine, 1,13 g (6 mmol) of *tert*-butyl 8-benzyl-8-azabicyclo[3.2.1]oct-3-yl-*exo*-carbamate and 0,97 ml (6,5 mmol) of 1,8-diazabicyclo[5.4.0]undec-7-ene were dissolved in 40 ml of *n*-pentanol and heated under reflux for 50 hours. The solvent was evapotared, the residue was dissolved in 50 ml of chloroform, washed with 4x30 ml of water, dried over sodium sulfate, and purified by column chromatography using *n*-hexane - ethyl acetate - chloroform (3:1:1) as eluent to result in white crystals which was triturated with *n*-hexane. Yield: 0,55 g (36 %). M.p.: 122-123°C. ¹H-NMR (DMSO-d₆): 8 1.34 (s, 9H), 1.44-1.66 (m; 2H), 1.67-1.99 (m, 6H), 3.88 (m, 1H), 4.56 (bs, 2H), 6.59 (d, 1H), 7.77 (d, 1H), 8.07 (dd, 1H), 8.17 (d, 1H).

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Example 4.

(2S)-1-(2-{[8-(5-Nitropyridin-2-yl)-8-azabicyclo[3.2.1]octan-3-yl]-exoamino}acetyl)pyrrolidine-2-carbonitrile

The meaning of R is 5-nitropyridin-2-yl group, B means a group of formula (1), Z means a group of formula (B) in general formula (I).

a.) <u>tert-Butyl 8-(5-nitropyridin-2-yl)-8-azabicyclo[3.2.1]oct-3-yl-exo-</u>

<u>carbamate</u> with (V) general formula, where R and B are given above, Y is tert
butoxycarbonyl group

476 mg (3 mmol) of 2-chloro-5-nitropyridine, 679 mg (3 mmol) of tert-butyl 8benzyl-8-azabicyclo[3.2.1]oct-3-yl-exo-carbamate and 0,46 ml (3,1 mmol) of 1,8-25 dissolved ml of diazabicyclo [5.4.0]undec-7-ene were in n-pentanol and heated under reflux for 1 hour. The solvent was evapotared, the residue was dissolved in 40 ml of chloroform, washed with 4x40 ml of water, dried over sodium sulfate and evaporated. The solid residue was triturated with diethyl ether to result in yellow crystals, Yield: 731 mg (70 %). M.p.: 212-214°C. ¹H-NMR (DMSO-d₆): δ 1.34 (s, 9H), 1.41-1.54 (m; 2H), 1.81-2.16 (m, 6H), 4.00 (m, 1H), 4.75 (bs, 2H), 6.63 (d, 1H), 6.82 (d, 1H), 8.21 (dd, 1H), 8.98 (d, 1H).

b.) 8-(5-Nitropyridin-2-yl)-8-azabicyclo[3.2.1]oct-3-yl-exo-amine with general formula (II), where R and B are given in step 4a.)

651 mg of tert-butyl 8-(5-nitropyridin-2-yl)-8-azabicyclo[3.2.1]oct-3-yl-exo-carbamate (1,87 mmol) was dissolved in 20 ml of 12% ethanolic hydrochloric acid and the solution was stirred for 3 hours. Under cooling 90 ml 1N sodium hydroxide was added to the formed a suspension which was extracted 4 x 50 ml dichloromethane. The layers were separated, the organic phase was dried, evaporated and the residue was triturated with n-